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Deep sequence analysis of non-small cell lung cancer: integrated analysis of gene expression, alternative splicing, and single nucleotide variations in lung adenocarcinomas with and without oncogenic KRAS mutations

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KRAS mutations are highly prevalent in non-small cell lung cancer (NSCLC), and tumors harboring these mutations tend to be aggressive and resistant to chemotherapy. We used next-generation sequencing technology to identify pathways that are specifically altered in lung tumors harboring a KRAS mutation. Paired-end RNA-sequencing of 15 primary lung adenocarcinoma tumors (8 harboring mutant KRAS and 7 with wild-type KRAS) were performed. Sequences were mapped to the human genome, and genomic features, including differentially expressed genes, alternate splicing isoforms and single nucleotide variants, were determined for tumors with and without KRAS mutation using a variety of computational methods. Network analysis was carried out on genes showing differential expression (374 genes), alternate splicing (259 genes), and SNV-related changes (65 genes) in NSCLC tumors harboring a KRAS mutation. Genes exhibiting two or more connections from the lung adenocarcinoma network were used to carry out integrated pathway analysis. The most significant signaling pathways identified through this analysis were the NFκB, ERK1/2, and AKT pathways. A 27 gene mutant KRAS-specific sub network was extracted based on gene–gene connections from the integrated network, and interrogated for druggable targets. Our results confirm previous evidence that mutant KRAS tumors exhibit activated NFκB, ERK1/2, and AKT pathways and may be preferentially sensitive to target therapeutics toward these pathways. In addition, our analysis indicates novel, previously unappreciated links between mutant KRAS and the TNFR and PPARγ signaling pathways, suggesting that targeted PPARγ antagonists and TNFR inhibitors may be useful therapeutic strategies for treatment of mutant KRAS lung tumors. Our study is the first to integrate genomic features from RNA-Seq data from NSCLC and to define a first draft genomic landscape model that is unique to tumors with oncogenic KRAS mutations.

Keywords: transcriptome sequencing, RNA-Seq, KRAS mutation, NSCLC, bioinformatics, network analysis, data integration and computational methods

INTRODUCTION

The most common form of lung cancer is histologically defined as non-small cell lung cancer (NSCLC). Activating mutations in the KRAS oncogene are often found in NSCLC patients with smoking history (Eberhard et al., 2005; Pao et al., 2005b). The KRAS oncogene harbors activating mutations, especially in codons 12 or 13; and such mutations are prevalent in pancreatic cancer (Almoguera et al., 1988), leukemia, colorectal carcinomas (Andreyev et al., 1997), and about 20–30% of lung adenocarcinomas (Riely et al., 2009). Another prevalent oncogene in NSCLC

is the epidermal growth factor receptor (EGFR). EGFR kinase domain mutations have been established as valid predictors of therapeutic response to EGFR-targeted therapeutics such as the small molecule EGFR inhibitors gefitinib, erlotinib, and lapatinib and the EGFR antibody cetuximab. In contrast, the therapeutic significance of KRAS mutations in NSCLC remains unclear and no clinically useful KRAS inhibitors have been developed for management of NSCLC patients (Riely et al., 2009). In NSCLC, activating KRAS mutations are predominant and are mutually exclusive of mutations in EGFR. Studies indicate that lung adenocarcinoma

patients with *KRAS* mutations are associated with resistance to *EGFR* inhibitors (Eberhard et al., 2005; Pao et al., 2005a; Mareselli et al., 2007). The mechanisms that underlie such resistance are largely unknown, and there is a very pressing need to identify and exploit new molecular targets for management of patients with NSCLC tumors with *KRAS* mutations. Since oncogenic *KRAS* has proved to be difficult to target directly (Vojtek and Der, 1998; Shields et al., 2000), an alternative strategy is to identify signaling pathways that are activated downstream of mutant *KRAS* and to develop key nodal components of these pathways as therapeutic targets using next-generation sequencing technology.

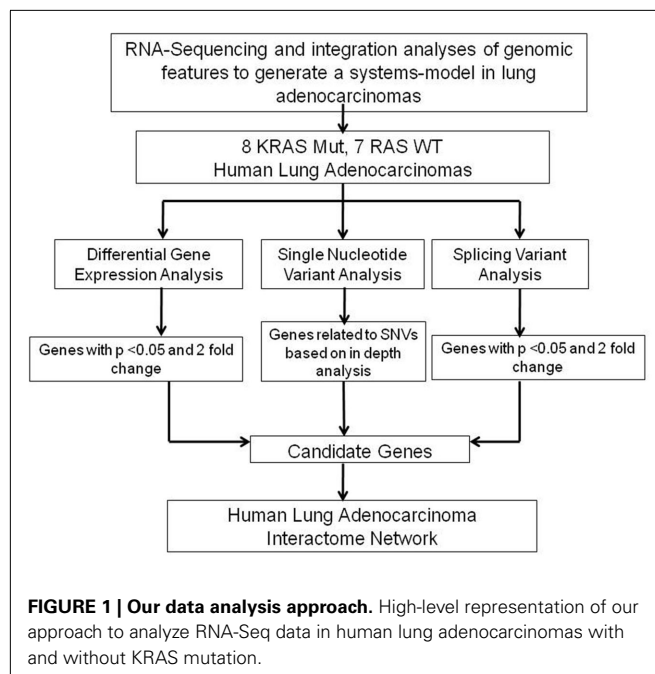
There is very little information on differential gene expression in NSCLC tumors with and without *KRAS* mutation. Interrogation of oncomine and gene expression omnibus (GEO) databases revealed few studies that have focused specifically on the relationship of *KRAS* mutation with gene expression in lung adenocarcinomas patients (Beer et al., 2002) or cell lines (Bild et al., 2006; Singh et al., 2009). Furthermore, most of these studies are based on Affymetrix Hu6800 oligonucleotide arrays and analytical technology that is, by modern standards, relatively immature to study gene expression profiles. Thorough analysis of microarrays led us to conclude that there is little reliable data on differential patterns of gene expression in NSCLC tumors with and without *KRAS* mutations, and virtually no genomic studies of somatic mutations, splice variants, or fusion gene products that are specifically associated with such tumors is available. Deep sequencing of transcriptome (RNA-Seq) provides a powerful tool to interrogate the whole transcriptional landscape. Therefore, we combined RNA-Seq with sophisticated methods and new analytical pipelines developed by our group to analyze RNA-Seq data, to revisit the challenge of identifying genomic features that define differences in the genomic landscape of *KRAS*-mutated and *KRAS*-wild-type primary NSCLC tumors.

In the present study, we identified genomic features such as differential gene expression, alternate splice variants, and expressed polymorphisms that are significantly involved in NSCLC tumors harboring *KRAS*-mutated tumors when compared to *KRAS*-wild-type tumors (Figure 1). These genomic features were then used to develop a human NSCLC interactome network. Our analysis represents the first reported effort to integrate gene expression, alternatively splicing, and nucleotide sequence variation data into a model that define a genomic landscape unique to NSCLC tumors harboring oncogenic *KRAS* mutations. Our results, in addition to validating previous studies on the role of RAF, ERK1/2, AKT, and NFκB in mutant *KRAS* NSCLC, also reveal novel links to other druggable target pathways including TNFR and PPARγ. Our results indicate that this approach will lead to novel insights into the biology of mutant *KRAS* tumors and identify novel druggable pathways to treat *KRAS*-mutant tumors.

MATERIALS AND METHODS

DATA SHARING

The sequence data used in this manuscript have been deposited in GEO (GSE34914).



SAMPLES

We performed RNA-sequencing of 15 lung adenocarcinomas, 8 with *KRAS* mutation and 7 without *KRAS* mutation. All tumors were grade I or II and were obtained from surgical resection. Tumors were macrodissected to remove normal tissue prior to freezing, and all samples were histologically evaluated and determined to be >70% tumor tissue. The *KRAS* mutational status was determined by polymerase chain reaction (PCR) amplification and confirmed by Sanger sequencing of exon 1 of *KRAS*. These studies were carried out under Mayo Clinic IRB protocol 08-005844.

RNA PREPARATION AND SEQUENCING

Total RNA was prepared from 15 fresh frozen lung adenocarcinomas. All RINs were >7.0, as determined using the Agilent Bioanalyzer. The cDNA libraries were prepared from polyA enriched RNA using Illumina protocols. cDNA fragments of 300–400 bp were selected, and non-directional 50 nucleotide paired-end sequencing was performed as described previously (Sun et al., 2011). Sequencing was carried out at Mayo Clinic Advanced Genomic Technology Center at Rochester, MN, USA using the Illumina Genome Analyzer II (GA II). One tumor sample without *KRAS* mutation was run twice for QC evaluation. The FASTQ read files for the 16 samples were used for further data analysis. Data for gene counts was obtained using our Mayo Clinic pipeline and Burrows–Wheeler Alignment (BWA) alignment. Twenty to fifty-two million tags were obtained from sequencing. The percent of reads mapped for 16 samples varied from 71 to 84.2%. Table 1 consists of details from sample statistics for paired-end runs; the table contains counts combined for each sample from both reads.

DATABASES AND SOFTWARE USED FOR ANALYSES

Gene expression and alternate splicing data analyses were carried out by downloading database tables from the UCSC table browser

Table 1 | Statistics based on per sample analysis using BWA alignment for paired-end reads.

Sample ID	LUS9	LU374	LU350	LU350	LU242	LU528	LU115	LU242_2	LU53	LU185	LU213	LU256	LU273	LU325	LU439	LU499
Total reads	20,439,292	35,911,708	36,093,910	35,563,802	35,556,342	36,584,220	38,580,592	34,563,748	39,682,456	41,166,162	41,513,982	42,064,264	40,663,636	41,746,522	40,413,428	39,981,816
Mapped reads	16,124,080	26,579,036	26,927,145	27,008,244	26,970,516	29,934,989	27,431,276	28,106,604	30,976,997	34,500,496	33,343,330	34,326,544	31,585,880	29,326,676	34,035,189	33,213,588
Percentage mapped reads	78.90	74.00	74.60	75.90	61.50	81.80	71.10	81.30	78.10	83.60	80.30	81.60	77.70	70.20	84.20	83.10
GeneCount	13,668,945	19,532,542	19,456,836	22,077,296	21,988,285	25,996,314	22,499,005	21,294,617	26,969,974	31,730,519	29,231,047	30,717,165	27,664,166	25,919,379	31,008,817	28,081,496
ReadStart	14,174,891	20,255,603	20,117,537	22,934,458	22,780,182	27,034,656	23,301,045	22,058,823	27,914,235	32,889,808	30,281,834	31,795,315	28,641,860	26,897,161	32,082,534	29,198,421
GeneCount ≥ 10 reads	15,871	15,754	15,841	16,043	16,335	16,087	15,287	16,249	15,915	15,936	15,709	16,609	16,347	15,741	16,071	15,318

in reference to human genome build GRCh37, which corresponds to UCSC hg19 assembly (Fujita et al., 2011). 1000g2010nov data was obtained from the 1000 Genomes Project PHASE, 2010 November release¹ and dbSNP version 132² was used for single nucleotide variation (SNV) analysis. SIFT database provided by <http://sift-dna.org/> was used to predict whether an SNV coding an amino acid substitution will affect protein function. ANNOVAR was used to functionally annotate genetic variants (Wang et al., 2010). TopHat (Trapnell et al., 2010) and BWA tools (Li and Durbin, 2009) were used to align RNA-Seq reads. Most of the statistical analyses were conducted using R: a language and environment for statistical computing³. Quantification of splicing from paired-end reads was performed using an R package – CASPER to infer gene alternative splicing patterns from paired-end sequencing data (Rossell, 2010). Partek software tools were used for plot generation and data mining purposes. A Microsoft SQL server database was used to store and query data for analysis. A series of computational programs was written using Perl scripting language to access and filter data from the Microsoft SQL database.

GENE EXPRESSION DATA ANALYSIS

The Illumina standard pipeline, GA II was employed for processing of raw images, to make base calls and to generate FASTQ sequence reads from paired-end RNA-sequencing data. The exon-exon boundary database was generated using exon and gene definitions obtained from UCSC refFlat table for hg19 assembly. Uni-directional combinations of exon junction database for the sequencing length (50 bases) were generated using exon boundaries defined by the refFlat file from UCSC Table Browser. FASTQ sequence reads were aligned to the human reference genome (hg19) and to our in-house exon junction database using BWA. BWA is a fast and accurate short read aligner. A maximum of two mismatches were allowed for first 32 bases in each alignment, and reads that had more than two mismatches or were mapped to multiple genomic locations (alignment score less than 3) were discarded. The aligned sequence tags were summarized and annotated using SnowShoes, an in-house RNA-Seq pipeline (manuscript in preparation). The read counts for genes are generated for further downstream analyses. A non-mutant KRAS sample that was run twice had high correlation; hence we took the average read count for each gene for that sample. Raw read counts for a total of 22,316 genes were obtained for gene expression analysis. Genes (15,092) that had a median read count $>2^4$ (16 reads) in at least one of the KRAS groups were used for further analysis. Individual read count data were normalized using mode, as follows: read count/sample read count mode *sm, where sm is the smallest mode across all the samples. Since the length of the transcript is assumed to be constant when comparing two sample cohorts, no normalization for the length of genes was performed for differential expression analysis. ANOVA implementation of Partek Genomics Software was used for differential gene expression analysis after normalizing RNA-Seq gene count data. Microarray data used in this study were normalized with gc-robust

¹<http://www.1000genomes.org>

²<http://www.ncbi.nlm.nih.gov/projects/SNP/index.html>

³<http://www.r-project.org>

multi-array average (gcRMA) algorithm, using Partek Genomics (Partek Inc., St Louis, MO, USA). Normalized microarray data were analyzed using ANOVA implementation of Partek Genomic Software for differential gene expression analysis.

ALTERNATE SPLICING DATA ANALYSIS

TopHat is a fast splice junction mapping software that uses short read aligner – Bowtie to align RNA-Seq reads (Langmead et al., 2009; Trapnell et al., 2010). TopHat outputs alignments in sequence alignment/map (SAM) format. Samtools (Li et al., 2009) were used to convert files to binary alignment/map (BAM) format. BAM files were loaded into Bioconductor, and the CASPER package⁴ was used to quantify known splicing variants. Briefly, CASPER obtains the list of known splicing variants for each gene from UCSC, and estimates their relative abundances by modeling the reads as a mixture of multinomial distributions (Figure 2). Maximum likelihood estimates of the relative abundances were obtained via the EM algorithm (Dempster et al., 1977). Transcript expression data was obtained for each sample from CASPER analysis. Transcripts with no mapped reads were imputed with a zero and all samples with transcript data were merged for splicing analysis. Mann–Whitney–Wilcoxin test that does not assume normality was performed to identify differential alternative splice forms between the *KRAS*-mutant and *KRAS*-wild-type samples.

SNV DATA ANALYSIS

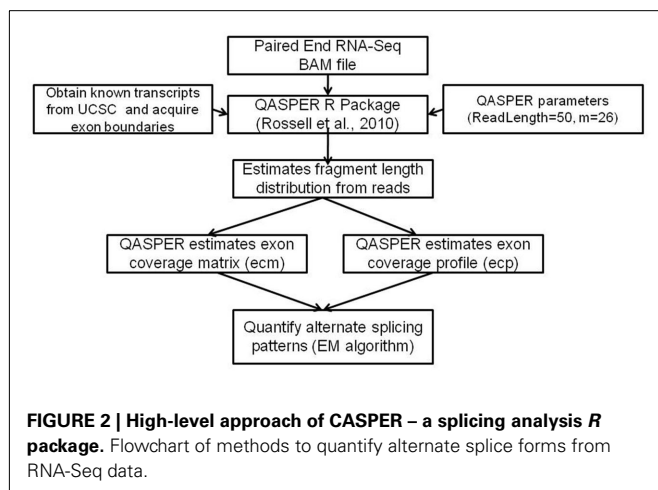
Novel SNVs and SNVs with different allelic frequencies in samples with and without *KRAS* mutations were also discovered using transcriptome sequencing data. Analysis of SNVs from lung adenocarcinomas has been performed using a variety of computational methods. Single reads from paired-end data for a sample were aligned to exon–exon junction database and genome independently with BWA default parameters (Figure 3). When a read maps to junction and genome, the read that map to the junction gets priority and the read from genome mapping will be filtered. In addition to duplicate mapping reads, reads of low quality (mapQ < 20) were also filtered. From a paired-end RNA-Seq, four files were obtained from junction and genomic BWA

alignments. As shown in Figure 3, a total of four BAM alignment files for each sample were used to create a pileup file. The pile up file generated for each sample was used to predict single nucleotide variants using SNVMix – a novel binomial mixture model (Goya et al., 2010). The number of reference and alternate base reads were obtained for every transcribed position in the genome. SNVMix uses a probabilistic method based on a binomial mixture model to infer genotypes. At each nucleotide position in the data, there is one of three genotype states (Goya et al., 2010) generated from sequence data: aa (homozygous for the reference base), ab (heterozygous), and bb (homozygous for the non-reference base). In order to minimize errors, only bases with >Q20 base quality were considered in determining counts. We also filtered our SNV data against dbSNP version 132, the 1000 genomes project (see text footnote 1), the Illumina Body Map 2.0 data from 16 normal tissues (data downloaded from www.broadinstitute.org/igvdata/BodyMap/hg19/IlluminaHiSeq2000_BodySites), and four normal lung epithelial cell samples (Beane et al., 2011). Illumina body map data and normal lung epithelial RNA-Seq was obtained in order to eliminate any previously described germ line variants. Illumina body map samples and lung epithelial cell lines were processed for SNVs in the same manner as our 16 lung adenocarcinoma samples. SNV data from all samples were stored in a Microsoft SQL server 2005 database. In-house programs were developed to filter and query data from the database. ANNOVAR software was used to functionally annotate and filter genetic variants. Frequency comparisons of the SNVs were performed using the Fishers exact test from Plink software⁵. Additional filtering of SNVs was performed using SNV data generated from TopHat alignment in order to remove any false positive SNVs arising due to alignment issues. A variety of filters were built on forward and reverse directionality reads for SNVs, as described in the text.

PATHWAY AND NETWORK ANALYSES

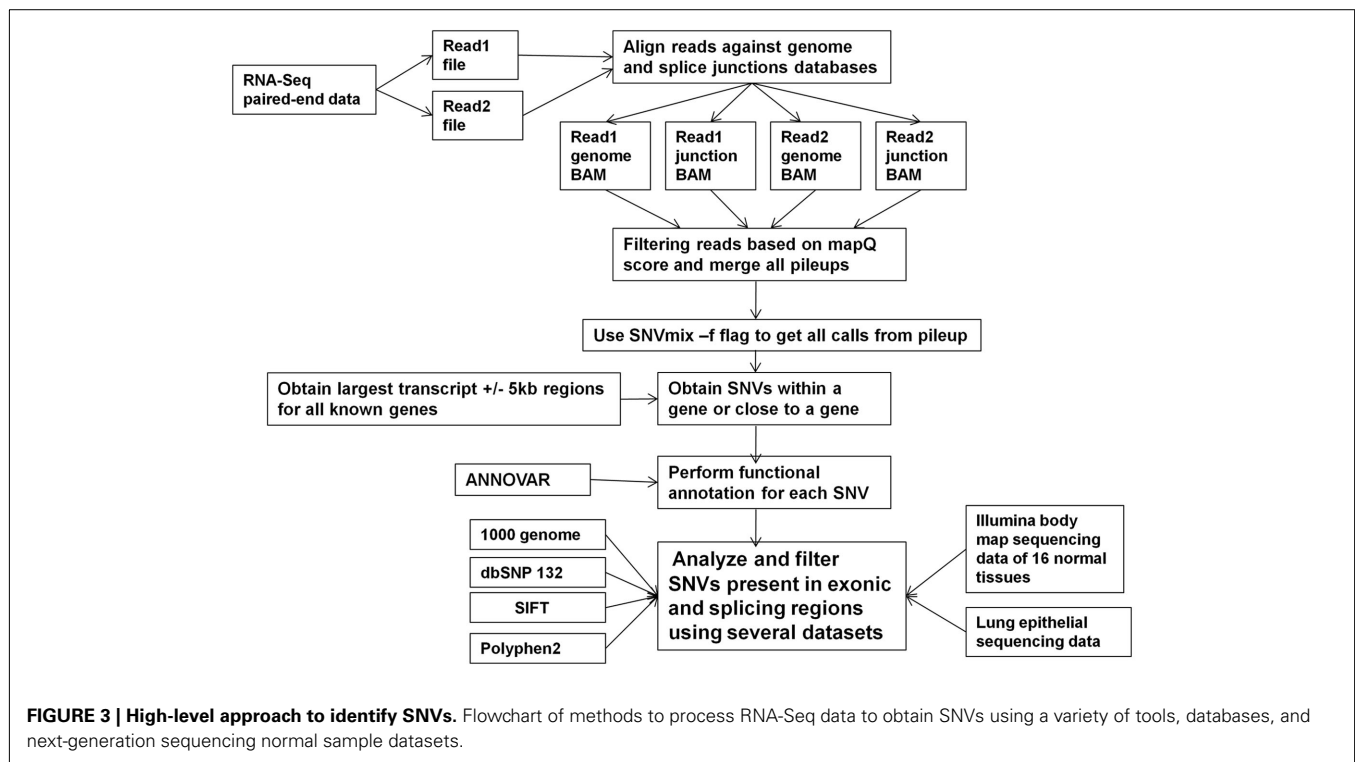
Ingenuity Pathway Analysis (IPA) software (Ingenuity® Systems⁶) was used for pathway analysis. IPA constructs protein interactions based on a regularly updated “Ingenuity Pathways Knowledge Database.” The IPA knowledge database consists of millions of relationships between proteins extracted from the biological literature. Each relationship between molecules is created using the IPA knowledge database. IPA was used to identify pathways based on differential gene expression, alternate splicing and SNV gene sets. Significant pathways in IPA were identified using Fisher’s exact test. The *p*-value indicates the likelihood of the input gene list or proteins in a given pathway or network being found together due to random chance. Cytoscape, a popular open source bioinformatics package, was used for complex network analysis and visualization (Smoot et al., 2011). Integration of the genomic features (differential expression, alternative splice variants, and SNVs) was performed using Cytoscape. Network analyzer, a Cytoscape plug-in, was used to compute a comprehensive set of topological parameters for directed networks (Assenov et al., 2008). Number of

⁴ <https://sites.google.com/site/rosselldavid/software>



⁵ <http://pngu.mgh.harvard.edu/~purcell/plink>

⁶ <http://www.ingenuity.com>



edges, distribution of degree counts, and neighborhood connectivity scores were obtained for the integrated network using network analyzer. Reactome was also used for network analysis. Reactome consists of expert curated and peer-reviewed high quality data to infer human functional interactions among genes⁷. Reactome and human interactome⁸ databases were also used along with Cytoscape to build networks. Only the genes that were expressed in RNA-Seq data (median read count > 16 in at least one of the *KRAS* groups) were used as reference gene set during network analysis.

RT-PCR AND SANGER SEQUENCING VALIDATION

Genomic DNA was extracted from NSCLC tumors using standard protocols. The seven SNVs corresponding to genes (*KRAS*, *GSTZ1*, *ECT2*, *GLS*, *WDTCL1*, *SRSF3*, and *RBM23*) were evaluated using Sanger sequencing. Primer pairs for these SNVs were designed with Primer3 version 4.0 software, and were used to amplify all variants by PCR. PCR products were purified from unincorporated nucleotides using a Millipore PCR purification plate. A total volume of 10 μ l, containing 80 ng of the clean product and 1.6 pM of one of the primers (forward or reverse), was used for sequencing. Electropherograms were analyzed with SeqScape v2.5 (ABI, Applied Biosystems, Foster City, CA, USA).

Quantitative real time PCR (qPCR) was used to verify alternative splicing of selected candidate transcripts. Total 1 μ g of RNA was isolated and converted to cDNA using Applied Biosystem's High Capacity cDNA RT kit (Part # 4368813). Note that the protocol for cDNA library construction for qPCR was different from that used for RNA-Seq analysis, so as to minimize potential

artifacts that might arise during cDNA conversion. Probes and primers corresponding to known exon/exon junctions were purchased from Life Tech (Applied Biosystems) and qPCR was carried out using an AB 7900HT analyzer.

RESULTS

DIFFERENTIAL GENE EXPRESSION ANALYSIS

Read counts were obtained for 22,316 genes from 15 lung adenocarcinoma samples with and without *KRAS* mutation. The read count data were normalized using mode normalization and log2 transformation as described in the Section "Materials and Methods." Genes with a median read count < 16 in lung adenocarcinoma samples with and without *KRAS* mutation were eliminated from further analysis. The remaining 15,092 transcripts were analyzed using ANOVA to identify 374 transcripts that were differentially expressed genes in *KRAS* mutation samples versus *KRAS*-wild-type samples with p -value < 0.05 and twofold-changes in gene counts. The association between the *KRAS* genotype and the top six differentially expressed genes is shown in **Figure 4A**. Hierarchical clustering of the 374 differentially expressed genes is shown in **Figure 4B**, in which total gene counts were standardized by the mean value among the samples.

Ingenuity pathway analysis was used to determine biological relationships among the 374 differentially expressed genes. The top five networks, based on Fisher's exact test, were associated with immunological disease, cell signaling, cell death, nervous system development, and function and cellular development pathways. The key nodes of the five networks are NF κ B, ERK1/2, AKT, MAPK, PI3K complex, IL12, and JNK. The top four networks (gene-gene relationships) from IPA analysis are shown in **Figures 5A–D**. IPA also determines the number of subgroups

⁷ <http://www.reactome.org>

⁸ http://cytoscape.wodaklab.org/wiki/Data_Sets

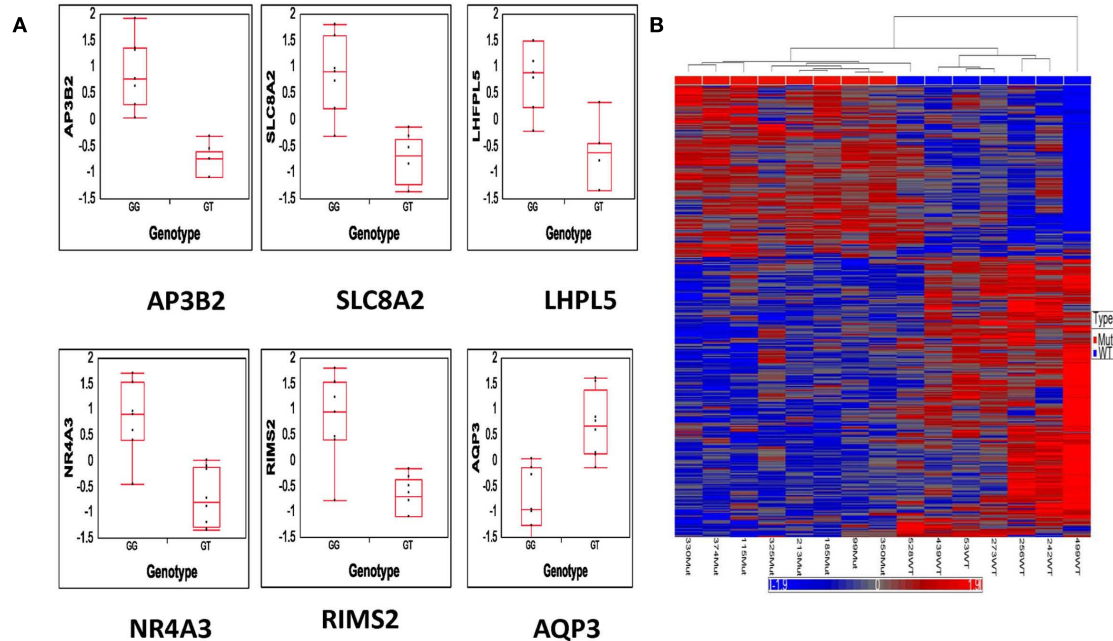


FIGURE 4 | Results from differential gene expression analysis. (A) Top six significantly differentially expressed genes between *KRAS*-mutant genotype (GT) and *KRAS*-wild-type genotype (GG). **(B)** Hierarchical clustering of 374 genes that are differentially expressed with at least twofold-change and p -value < 0.05 .

of genes that are associated with a known or canonical pathway. The top 3 canonical pathways from the 374 gene list are cell cycle regulation by B-cell translocation gene (BTG) family proteins, glioblastoma multiforme signaling, and Wnt/ β -catenin signaling.

ALTERNATIVE SPLICING ANALYSIS

We used CASPER, an R package, to obtain the expression of isoforms for all the refseq genes or transcripts. The UCSC hg19 assembly consists of 31,599 known transcripts. Genes with only one known isoform were removed from the analysis (13,633 transcripts). CASPER estimates relative expression levels, i.e., proportion of transcripts for a given gene originating from each variant. Transcript expression values from individual samples were organized into a single file and transcripts with no reads were set to zero. Data from CASPER files contained 17,966 transcripts for further analysis. There were 314 transcripts corresponding to 259 genes with a p -value < 0.05 and minimum twofold-change in median of multiple ratios between *KRAS*-mutant and wild-type samples. An example of output from CASPER package is shown in **Figure 6**. The estimated abundance of alternatively spliced NM_00268 and NM_053024 transcripts is 66.79 and 33.21 for the profiling two gene *PFN2* in *KRAS* mutated samples (**Figure 6A**), whereas the abundance ratios for NM_00268 and NM_053024 transcripts are estimated as 82.28 and 17.72% respectively for *KRAS*-wild-type samples (**Figure 6C**). **Figures 6B,D** show the reads for *PFN2* transcripts for the same *KRAS* mutated and wild-type lung adenocarcinoma samples. We also analyzed data using junction data obtained from TopHat to identify isoforms. Our *in silico* analysis of junction data for *PFN2* gene from TopHat is similar to CASPER data and indicates that the short isoform (NM_002628)

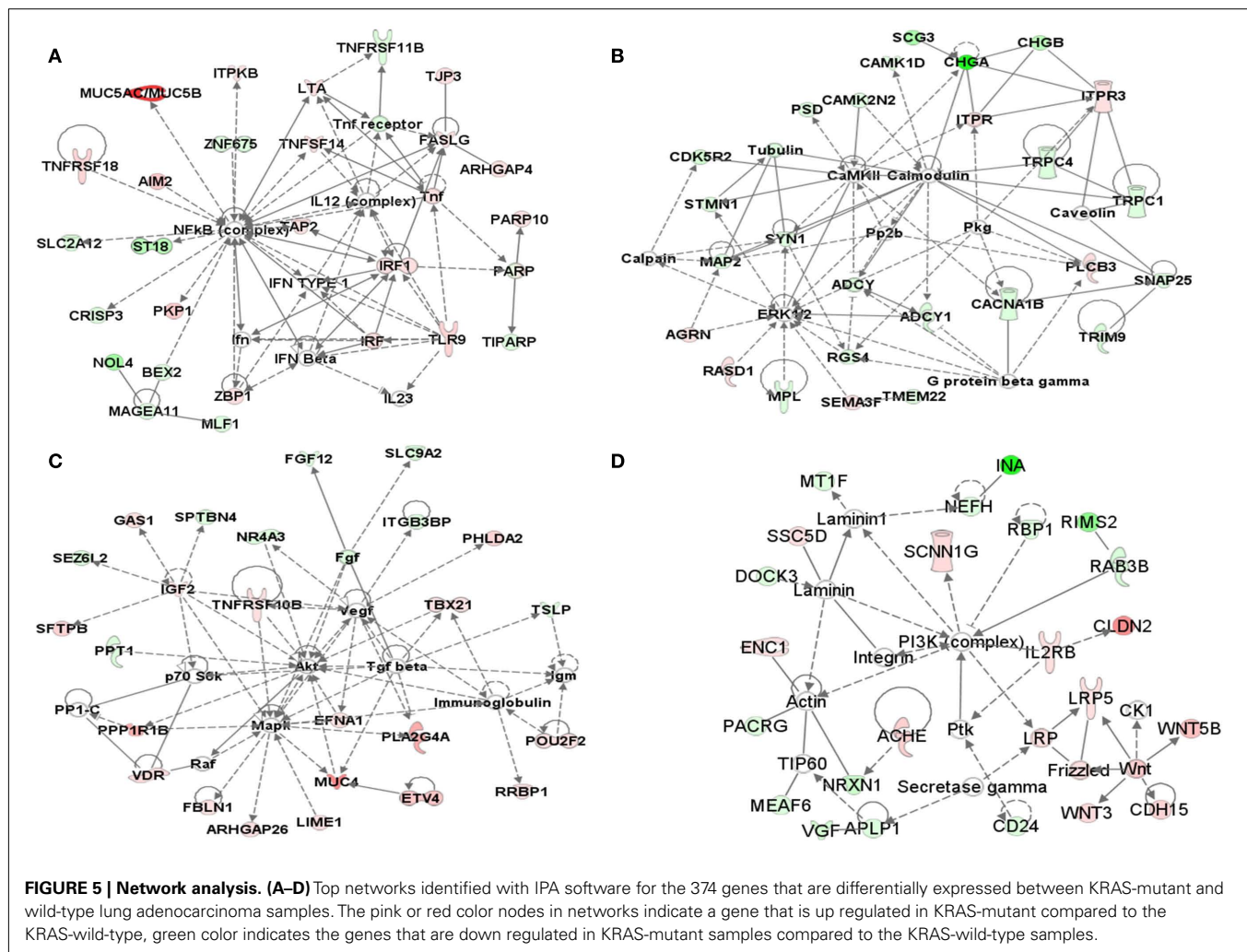
is abundantly expressed in *KRAS*-mutant samples compared to *KRAS* WT group.

For pathway analysis using IPA, we have used the 259 genes corresponding to 314 transcripts. The top five networks from IPA analysis were associated with cellular growth and proliferation, cell-to-cell signaling and interaction, nervous system development and function, molecular transport, and infection mechanisms. Similar to the pathway analysis of differentially expressed genes, the key nodes in the top five networks were NF κ B, ERK1/2, SNCA, AKT, PKC, MAPK, and Insulin.

SNV ANALYSIS

The SNVs that are ± 5 kb from a refseq gene were obtained and filtered based on several criteria as described in Section “Materials and Methods.” From the SNVMix output, genotypes ranging from 60 to 120 million per sample were investigated to call an SNV.

Nucleotide sequence at every genomic coordinate was evaluated when we had sufficient depth of sequence. We evaluated each nucleotide position for read depth, and for reads supporting the reference and alternate alleles for an SNV. An SNV was called when the genotype of one or more samples deviated from the reference genome genotype. In the current analysis, a SNV was not investigated if the genotype consists of multiple alleles, if there was no variation in genotype, or if the variation is present in less than two samples. We also eliminated SNVs from further analysis if the read depth was < 3 . After the application of the filters described above, a final set of 73,717 unique single nucleotide variants remained for further investigation. Since we have considered the regions that are close to a gene for SNV investigation, the majority of the SNVs are present in exonic (34,411) and 3' UTR (25,736) regions. Of the 34,411 exonic SNVs, there are 11,949 synonymous SNVs that



do not change an amino acid. Hence, we ignored these SNVs and estimated alternate allele frequencies from genotype data obtained from SNVMix software between the lung adenocarcinomas groups in 23,987 non-synonymous, exonic SNVs. For additional investigation of SNVs, we also obtained the 1000 genome and dbSNP132 data for all the 23,987 SNVs which include non-synonymous, stop gain, and stop loss mutations.

Fisher's exact test, called from PLINK software, was used to calculate differences in alternate allele frequencies. Eighty-four SNVs corresponding to 74 genes had a p -value <0.05 between tumor samples with and without *KRAS* mutation. **Table 2** shows details of 13 SNVs with a p -value <0.01 . **Table 2** consists of 18 columns with a variety of information for 13 SNVs.

Table 2 consists of details of SNV (chr, chromosome; Position, chromosome location; Gene, gene corresponding to SNV; Ref, reference allele; Alt, alternate allele; *KRAS* Mut, frequency of alternate allele observed in *KRAS* mutation group; *KRAS* WT, frequency of alternate allele observed in group without *KRAS* mutation; MAf, minor allele frequency; P , Fishers exact p -value; 1KGenome, frequency of alternate allele observed in 1000 genome samples; dbSNP132, dbSNP ID based on chromosome location if it exists;

SIFT, SIFT score that predicts amino acid changes that may be affected by protein function; MAF Body Map, frequency of alternate allele in Illumina body map 16 tissues that are sequenced and analyzed similarly for SNVs; Illumina Lung, genotype of Illumina lung sample from body map data and lung epithelial genotypes for NormNonSmoker – normal non-smoker; NormSmoker, normal smoker; SmokerNon-Cancer, smoker non-cancer; and SmokerLungCancer, smoker lung cancer samples respectively). “?” represents data not available in **Table 2**.

As shown in **Table 2**, the top significant SNV located on chromosome 12 corresponded to the *KRAS* gene. Thus, our SNV analysis confirms our *in silico* validation of sample classification based on *KRAS* mutational status. In addition to *KRAS* mutation there are an additional 79 SNVs that are present in the 1000 genome dataset with a minor allele frequency (maf) ranging from 0.95 to 0.02 (median maf = 0.35) and four more SNVs that are not present in the 1000 genome or dbSNP132 datasets, but which are differentially observed in the *KRAS*-mutant and wild-type samples with Fisher's exact p -value <0.05 .

To test if any of the four SNVs, corresponding to *WDC1* (chr1:27630115), *GLS* (chr2:191819311), *SRSF3* (chr6:36564670),

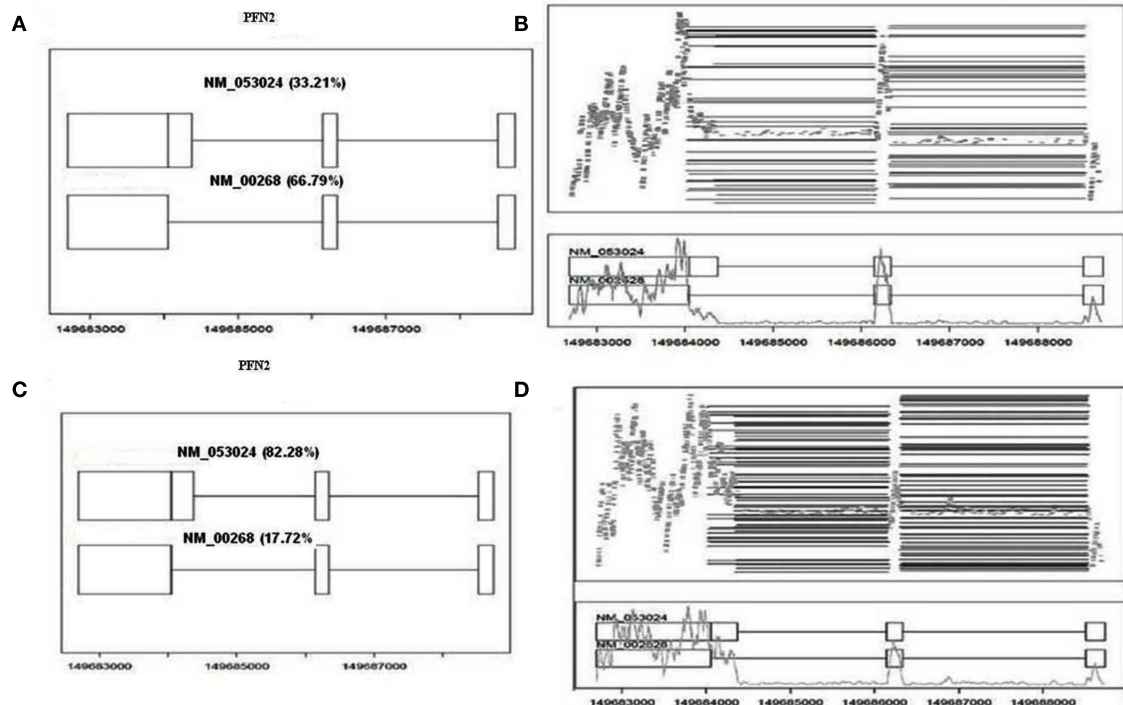


FIGURE 6 | Output of CASPER from splicing analysis. (A) CASPER output for alternate splice forms in *KRAS*-mutant sample. The data indicates that NM_00268 transcript is highly abundant compared to NM_053024 transcript of *PFN2* gene. **(B)** Observed reads for *PFN2* gene. Black lines in upper section of the plot indicates long reads and gray lines

indicate short reads typically spanning a single exon. The gray line on bottom half of plot indicates coverage at each position. **(C)** CASPER output for *KRAS* WT sample. NM_053024 transcript is predominantly expressed compared to other transcript. **(D)** Is similar to **(B)** but it corresponds to *KRAS* WT sample in **(C)**.

and *RBM23* (chr14:23370943) are novel, we performed TopHat alignment and individually examined the data in nucleotide sequence pileup files created from TopHat bam files. SNVs corresponding to *RBM23* and *GLS* are located at exonic splicing junction. Hence the alignments for these two variants were also examined carefully in integrative genomic browser (IGV). We found that the reads supporting the alternate allele for these two variants were present at the 5' ends of the sequence reads, and therefore likely to represent sequence and/or alignment errors. Hence we dropped these two SNVs from our analysis. The SNV corresponding to *WDTC1* was not called as a variant during TopHat alignment. There were no reads supporting the alternate allele, hence we discarded that as a novel variant. Finally, we investigated the variant corresponding to *SRSF3* from TopHat alignment and pileup data. This variant had an average read depth of 184; however, we found there is a strand bias of reads for the alternate allele. Thus, reads supporting the alternate allele C in the forward strand were 21, compared to 565 reads from the negative strand, such that this SNV was also eliminated on this basis.

We then filtered the remaining differentially detected SNVs to eliminate those that are at a splice junction, have alternate alleles that are exclusively found at the 5' end of the reads, or have strand bias for alternate reads. These rules plus a limit for maximum read depth to call an SNV and genotype confidently using SNVMix were applied to remove false positives. This led us to 72 SNVs

corresponding to 65 genes that have different allelic frequencies between the tumors with and without *KRAS* mutation.

An IPA analysis was carried out on the 65 genes corresponding to SNVs that have a preferential alternate allele present in samples with or without *KRAS* mutation. The top five networks were associated with tumor morphology, cellular growth and proliferation, cell death, cellular function and maintenance of cancer, neurological disease, cellular development, cell cycle, and cell morphology. The most significant disease associated with these SNVs was cancer and the key nodes in the top five networks were NFκB, ERK1/2, AKT, TNF, PI3K, ESR1, beta estradiol, and TGFβ1.

INTEGRATION ANALYSIS

Genes from differential expression (374 genes), alternate splicing (259 genes), and SNV (65 genes) analyses were used for integration analyses. Genes existing in multiple features were removed such that the final gene set for integration analysis consisted of 659 unique genes. Chromosomal mapping of these 659 genes identified several lung adenocarcinoma-specific clusters (Figure 7). Individual evaluation of chromosomes suggest that there may be clusters of genes on chromosomes 1, 3, 6, and 11 that are associated with *KRAS* lung adenocarcinomas (Figure 7).

NFκB, ERK1/2, and AKT canonical pathways were observed in all three independent feature analyses and also in the integrated analysis of 659 genes. The NFκB pathway has previously been

Table 2 | Top 13 SNVs that have different allelic frequencies in KRAS mutation group compared to the KRAS-wild-type samples.

Chr	Position	Gene	Ref	Alt	KRAS Mut	KRAS WT	MAF	P	1KGenome	dbSNP 132	SIFT	MAF body map	Illumina lung	Norm non-smoker	Norm smoker	Smoker non-cancer	Smoker lung cancer
12	253982S5	KRAS	C	A	8	0	0.3	0.002	?		0.03	0	CC	CC	CC	?	CC
14	77793207	GSTZ1	G	A	8	0	0.3	0.002	0.34	rs7975	0.04	0.313	AA	GA	GA	AA	GA
22	19951271	COMT	G	A	2	11	0.4	0.003	0.38	rs4680	0.09	0.469	GA	GA	GA	GA	GA
16	427479	TMEM8A	T	C	12	3	0.5	0.004	0.51	rs11248931	?	0.469	TC	TC	CC	CC	CC
16	426432	TMEM8A	T	C	12	3	0.5	0.004	0.49	rs2071915	1	0.469	TC	TC	TC	TC	CC
17	73552185	LLGL2	G	A	12	3	0.5	0.004	0.39	rs1671036	0.04	0.464	GA	GG	GG	GA	GA
6	31324200	HLA-B	G	C	9	1	0.3	0.006	0.62	rs1140412	1	NA	CC	GC	GG	GG	GG
11	111896324	DLAT	C	T	1	9	0.3	0.006	0.35	rs2303436	0.23	0.281	CC	CT	TT	CC	CC
16	69745145	NQO1	G	A	7	0	0.2	0.007	0.24	rs1800565	0.02	0.156	GG	GG	GG	GA	GG
19	44156472	PLAUR	T	C	7	0	0.2	0.007	0.14	rs2302524	0.03	0	TT	TT	TT	TT	TT
7	45104131	CCM2	G	A	7	0	0.2	0.007	0.12	rs11552377	0.61	0.125	GG	GG	GA	GG	GG
7	77247821	PTPN12	G	A	10	2	0.4	0.009	0.77	rs9640663	0.35	0.313	GA	AA	7	?	GG
16	30536918	ZNF768	C	G	10	2	0.4	0.009	0.3	rs10871453	?	0.313	CC	CG	CG	CC	CG

shown to be essential for the development of tumors with *KRAS* mutation in a mouse model of lung adenocarcinoma (Meylan et al., 2009). Previous studies have also shown the involvement of AKT (Okudela et al., 2004) and ERK1/2 canonical pathways (Blasco et al., 2011) in mutant *KRAS* lung adenocarcinomas. In independent feature analyses, the MAPK pathway is targeted by both differential gene expression and alternate splicing but not with SNVs. Similarly, PI3K has been shown to target through differential gene expression and SNV features but not by alternative splicing. These data indicate pathways involved in mutant *KRAS* tumors are targeted by multiple genetic mechanisms in these tumors.

RT-PCR GENE VALIDATION

We randomly selected 6/15 samples for qPCR validation of four differentially expressed genes identified in our RNA-Seq analysis. Significant correlations were observed ranging from 0.69 to 0.84, when qRT-PCR results were compared with RNA-seq expression data (data not shown). We also randomly selected two small genes with multiple splice variants for functional validation.

Two genes, one with two splice variants and another with more than two splice variants were used for functional validation of our splicing analysis. As shown in **Figure 8**, qPCR data of samples agree with CASPER analysis for *PFN2*. *KRAS*-mutant samples preferentially express the NM_002628 variant of *PFN2* when compared to the NM_053024 variants (**Figure 8**). The two most significant SNVs (*KRAS* and *GSTZ1*) showing differences in frequency in *KRAS*-mutant versus wild-type tumors were validated using Sanger sequencing of genomic DNA. The presence of the reference and alternate allele (A) is shown in IGV Browser (**Figure 8B**). Sanger sequencing validation of the SNV is also shown in **Figure 8C**.

NETWORK ANALYSIS

A 659 gene set obtained from our integration analysis was used to build lung adenocarcinoma networks. Reactome and protein-protein interactome databases were used to build networks using Cytoscape. Linker genes (genes not present in our list but known to interact with genes in our list based on the published literature or protein-protein interaction databases) were also included to construct networks using Cytoscape. Thus, our lung adenocarcinoma network consists of linker genes and the set of genes obtained from different genomic features of RNA sequencing analyses. The functionality and directionality of connections between the nodes are also indicated in **Figure 9**. Topological parameters of the directed network were obtained using network analyzer. **Table 3** consists of the top 50 genes from this analysis along with edge count, degree, and neighborhood connectivity parameters. As shown in the **Table 3**, *MAPK14* is a linker gene consisting of 63 edge counts, 12 outgoing edges, and 51 in degree edges, with a high neighborhood connectivity score of 23.71. Most of top genes from network analysis are linker genes, due to the fact that most of these genes are well established regulators of our candidate genes derived from different features of tumors. In summary, the most significant connected genes (hubs) from our analyses are *MAPK14* (linker gene), and *CCND1* from differential gene expression, *LAMA4* from SNV, and *RPS27A* from alternate splicing analysis.

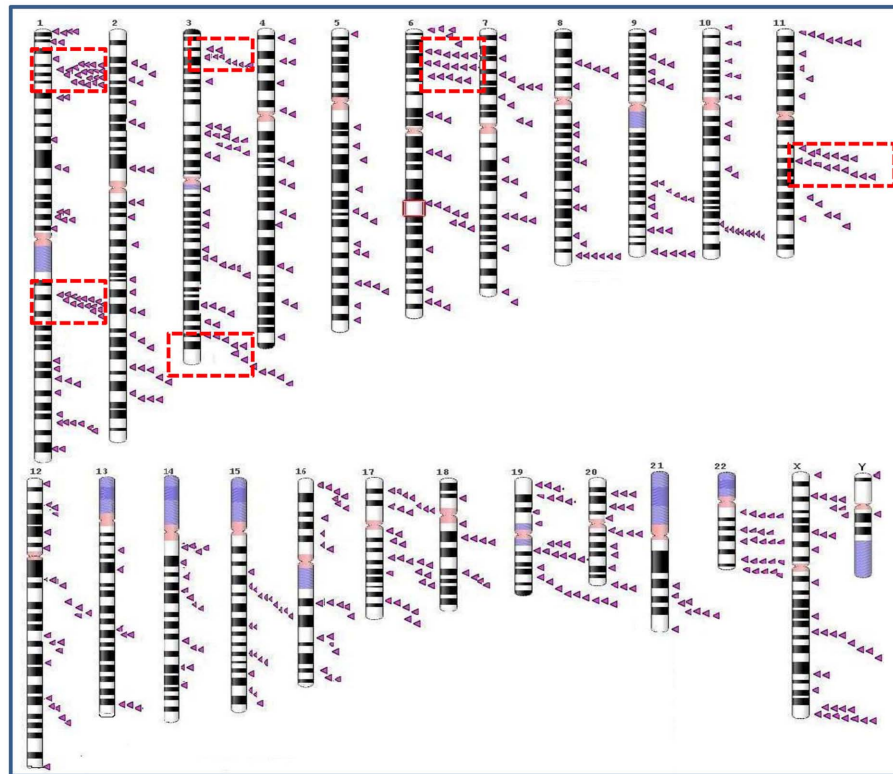


FIGURE 7 | Genomic view diagram. Chromosomal view of all the genes obtained from multi-feature analysis of lung adenocarcinomas with and without *KRAS* mutation. Chromosomes 1, 3, 6, and 11 consists of abundant gene clusters associated with *KRAS* mutation. Arrow in the diagram

represents a gene obtained from genomic feature analysis. Arrows identify the loci of genes obtained from genomic feature analysis. Larger arrows are shown when the genes are far apart, but when the gene locations are adjacent they are represented as smaller arrows.

To understand the connections between these genes in terms of known druggable targets and pathways we used IPA software. We obtained 387 genes that had at least two or more edges from the lung adenocarcinoma network developed previously (**Figure 9**) and submitted these data to IPA analysis. The 387 gene set consisted of 171 linker genes and 216 genes that were obtained from the multi-feature analysis of lung adenocarcinomas with and without *KRAS* mutation. In this way, we identified canonical pathways that are prevalent in the 387 dataset. **Figure 10** represents the top canonical pathways identified in our IPA analysis. The most significant canonical pathway consisted of 52/377 genes associated with molecular mechanisms of cancer (**Figure 10**).

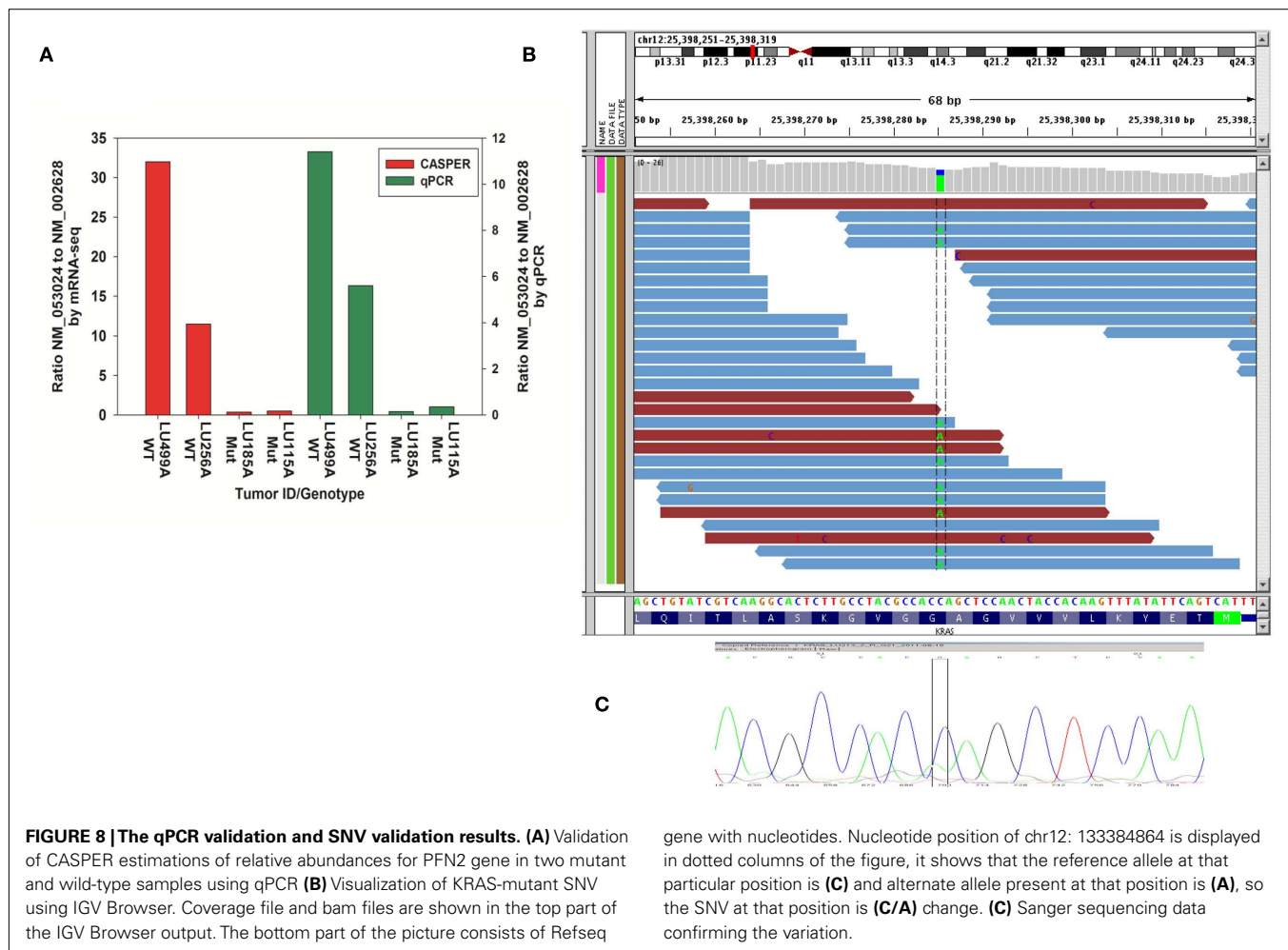
IDENTIFICATION OF KEY DRUGGABLE NODES FROM *KRAS* LUNG ADENOCARCINOMA NETWORK

To specifically explore the connections of the lung adenocarcinoma network with the *KRAS* gene, we obtained the upstream and downstream connections that are specific to *KRAS* gene from our 659 integrated multi-featured gene list. **Figure 11** shows the direct and indirect connections to *KRAS*. As shown in **Figure 11**, there are 14 genes (*LOC100506736*, *FBLN2*, *CCND1*, *AGRN*, *FBN1*, *MYCN*, *NQO1*, *SCNN1*, *ST5*, *TNFRSF10B*, *PPARG*, *GAS1*, *PLCB3*, and *IGF1*) that are indirectly connected with *KRAS* from our 659 unique gene list obtained from three different genomic features.

To build and expand the *KRAS* sub-specific network, we also used the grow feature in IPA to obtain direct gene-gene interactions from the Ingenuity Knowledge Base. This analysis gave us an additional 12 genes (*RAF1*, *RALGDS*, *RASSF1*, *ICMT*, *PTBP1*, *ELAVL1*, *WT1*, *ESX1*, *Ras*, *FEZF*, *RASGRF2*, and *IGF2BP1*) with direct connections to *KRAS*. The IPA overlay feature for known drugs was also used to determine if there are known FDA-approved drugs or clinical drug candidates within the 27 gene *KRAS* sub network. Three of the 27 genes are the target of known FDA-approved drugs (*RAF1*, *PPARG*, and *TNFRSF10B*).

In the 27 gene *KRAS* sub network we identified four known biomarkers for NSCLC (*CCND1*, *KRAS*, *PPARG*, and *Ras*). From the Ingenuity knowledge base, it is known that human *CCND1* protein and *PPARG* are useful biomarkers for measuring the efficacy of the PPAR γ agonist pioglitazone hydrochloride in the treatment of NSCLC. Similarly *KRAS* has been used as a biomarker for cetuximab, pazopanib, carboplatin, and erlotinib treatment of NSCLC. Interestingly, three therapeutic trials using *PPARG* agonists in NSCLC are currently being conducted⁹. Given our current analysis, it will be of interest to determine whether a correlation between clinical benefit and *KRAS* mutational status emerges from these trials.

⁹ Clinicaltrials.gov



DISCUSSION

We have conducted an integration analysis of multiple genomic features obtained from deep sequencing of NSCLC tumors with and without *KRAS* mutation. To our knowledge, this report represents the first attempt to leverage RNA-Seq data and a variety of computational methods to obtain an integrated genomic landscape map that incorporates differential gene expression, alternate splicing, and SNV data. This approach allowed us to comprehensively mine lung cancer data to produce a genomic landscape of NSCLC tumors. In our study, we took advantage of the genomic features obtained from our NSCLC samples along with the 1000 Genome data (normal samples), dbSNP data, and the Illumina body map data obtained from 16 normal tissues and the normal lung cell line data (Beane et al., 2011). Thus far there is only one paper that published lung cancer data using RNA-Seq; in that study, Beane and coauthors studied gene expression differences determined by RNA-Seq to understand the impact of tobacco smoke exposure in pooled samples from non-transformed bronchial epithelial cells from smokers and non-smokers with and without lung cancer. We used these data as “normal” controls for our analyses. To date, there is very little genomic information on differential gene expression, alternate splicing, and single nucleotide polymorphisms in NSCLC tumors with and without

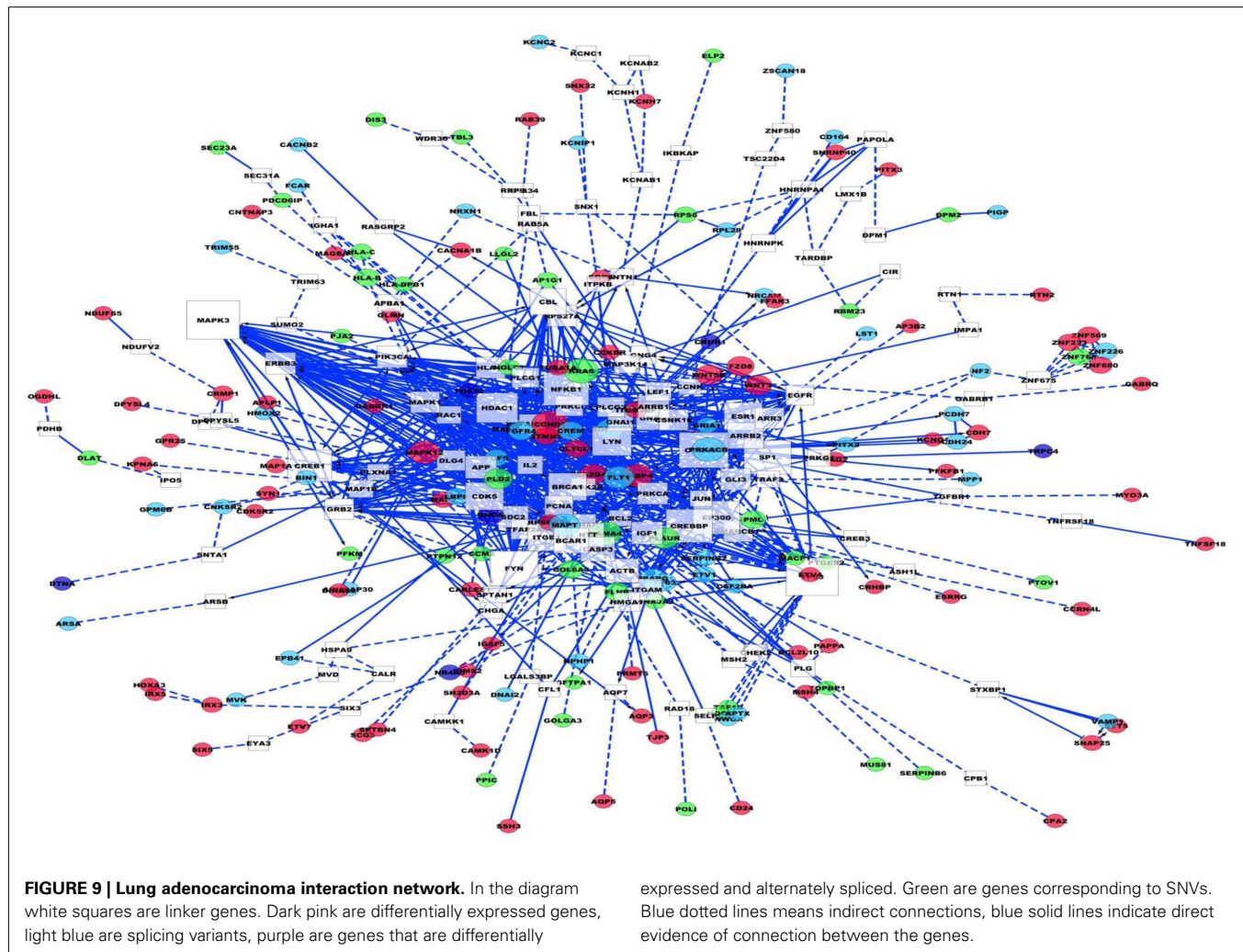
gene with nucleotides. Nucleotide position of chr12: 133384864 is displayed in dotted columns of the figure, it shows that the reference allele at that particular position is (C) and alternate allele present at that position is (A), so the SNV at that position is (C/A) change. (C) Sanger sequencing data confirming the variation.

KRAS mutation. We searched oncomine, GEO databases for gene expression studies and identified only two studies for which *KRAS* status of the tumors was known. One study was published in 2002 and used Affymetrix Hu6800 gene expression chips for 96 lung adenocarcinoma samples whose clinical parameters are known (Beer et al., 2002). Collected prior to 2001, the Beer et al. cohort contained 40 *KRAS*-mutants and 45 *KRAS*-wild-type samples, the remainder being of unknown *KRAS* status. Microarray data were reanalyzed to identify differentially expressed genes using the approach described in Section “Materials and Methods.” Only 19 genes were differentially expressed at p -value <0.05 and fold change $> \pm 2$. More than likely these results reflect analytical deficiencies related to the state of the art in microarray development that prevailed when these samples were analyzed. Another study determined *KRAS* status in 38 samples for human mammary epithelial cells and generated Affymetrix Human Genome U133 Plus2.0 (GSE3141) data (Bild et al., 2006). We reanalyzed GSE3141 data for which *KRAS* status was accurately published to identify differentially expressed genes using the approach described previously. Thirty-eight samples (11 samples with *KRAS* mutation and 27 samples without *KRAS* mutation) were used for analysis. Four hundred thirty-five probe sets corresponding to 311 genes were differentially expressed with a p -value <0.05 and with a fold

Table 3 | List of top 50 nodes from lung adeonocarcinoma network.

Gene	Edge count	Indegree	Outdegree	Neighborhood connectivity	Type
MAPK14	63	51	12	23.71	Linker
JUN	55	3	52	21.67	Linker
SP1	54	51	3	22.52	Linker
TP53	51	9	42	22.08	Linker
EP300	51	9	42	21.51	Linker
CREBBP	50	23	27	21.54	Linker
FYN	50	31	19	19.46	Linker
GRB2	50	34	16	22.26	Linker
CTNNB1	49	0	49	19.55	Linker
EGFR	46	16	30	25.76	Linker
NFKB1	45	28	17	23.56	Linker
RAC1	44	6	38	18.52	Linker
PIK3CA	43	0	43	27.12	Linker
MAPK3	43	5	38	26.30	Linker
AKT1	43	33	5	29.40	Linker
STAT3	41	29	12	26.49	Linker
CREB1	40	21	19	23.38	Linker
ITGB1	37	31	6	18.92	Linker
GNAI1	35	6	29	19.71	Linker
AR	35	6	29	28.97	Linker
RHOA	35	12	23	22.46	Linker
HDAC1	33	14	19	26.85	Linker
IL2	32	5	27	30.00	Linker
CBL	32	16	16	19.81	Linker
PRKACA	32	26	8	23.50	Linker
GNB1	30	10	20	20.23	Linker
EGR1	30	26	4	25.10	Linker
PRKCB	30	28	2	23.77	Linker
GNAO1	28	13	15	20.21	Linker
VEGFA	28	16	12	32.43	Linker
RXRA	26	1	25	24.35	Linker
RASA1	26	2	24	21.54	Linker
GNAQ	26	6	20	18.00	Linker
SP3	26	25	1	22.54	Linker
CALM1	25	19	6	18.12	Linker
ITGB3	25	20	5	22.00	Linker
HTT	25	21	4	24.96	Linker
LAMA4	24	16	8	21.33	SNV
CCND1	23	6	17	32.04	DE
ARRB2	23	6	17	20.17	Linker
TFAP2A	23	8	15	17.65	Linker
CASP3	23	9	14	21.13	Linker
ITGB2	23	18	5	18.17	Linker
RPS27A	22	12	10	21.14	AS
WT1	22	15	7.11	26.32	Linker
CRK	22	22	0	25.77	Linker
GNG2	21	1	20	15.95	Linker
ERBB2	21	3	18	28.19	Linker
GLI1	21	12	9	20.71	Linker
BRCA1	21	19	2	28.00	Linker

The statistics are based on the number of edge counts, neighborhood connectivity, in degree, out degree connection parameters obtained from network analyzer application in Cytoscape.



change $> \pm 2$. Only nine genes overlapped with our 345 differentially expressed genes. This could be due to differences in the gene expression patterns induced by oncogenic *KRAS* in breast versus lung tissue.

Deep sequence analysis of total mRNA makes it possible to analyze tumor samples and to quantify differential gene expression, alternative splicing, and SNV. There is always a concern that differences in cellularity may affect the outcome of genomic analyses of tumor tissue. We have attempted to minimize this potential problem by selecting only tumor samples that were histologically evaluated as $>70\%$ tumor tissue. In addition, there are several bioinformatics challenges such as base calling, assembly, alignment, and after-alignment handling of huge amounts of data from an RNA-Seq experiment. There are more than 300 software tools for next-generation sequence alignment, assembly, base calling, and post-alignment analysis¹⁰. We have used a combination of software tools for various analytical purposes. For each genetic feature obtained in this study, we computationally validated our analysis using at least two or more combinations of

approaches. We have discussed our approach for each genetic feature in detail in Section “Materials and Methods.” Our results demonstrate that transcriptome sequencing has the potential to provide new insights in our understanding of the genomic consequences of *KRAS* mutation in NSCLC patients by integrating different genomic features.

Even though *KRAS* mutations are highly prevalent in cancers, it has proven to be quite difficult to exploit mutated *KRAS* as a therapeutic target. Thus, a goal of our analysis was to identify druggable targets that are genomic associated with the mutant *KRAS* phenotype. Our analysis identified the *RAF*, *ERK1/2*, and *NFκB* pathways as specifically associated with mutant *KRAS*. Each of these pathways have previously been shown to be functionally linked to mutant *KRAS* tumorigenesis and represent bonafide drug targets for clinical treatment of *KRAS* tumors. Sorafenib inhibits *RAF1* and is currently being tested in NSCLC patients in phase II trial¹¹ (Blumenschein et al., 2009). In addition to these previously known *KRAS* related drug targets, our analysis revealed underappreciated links of mutant *KRAS* with the

¹⁰ <http://www.seqanswers.com>

¹¹ <http://www.clinicaltrials.gov/ct2/show/NCT00870532>

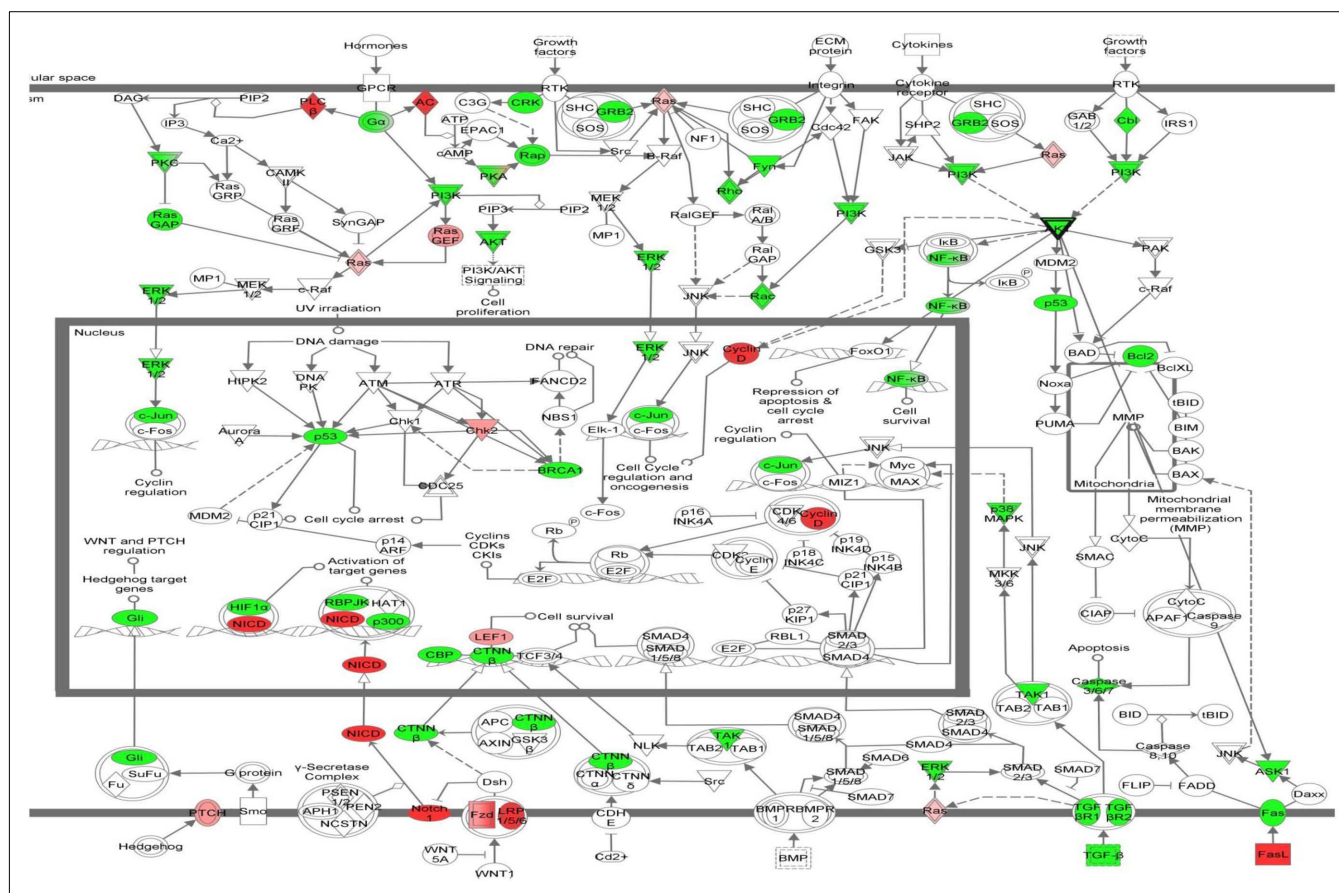


FIGURE 10 | Most significant canonical pathway from this study.

Molecular mechanisms of cancer is the most significant known pathway obtained from multi-feature analysis of lung adenocarcinoma

network. In the diagram green symbols represent linker genes and red or pink colors represent list of genes from lung adenocarcinoma network.

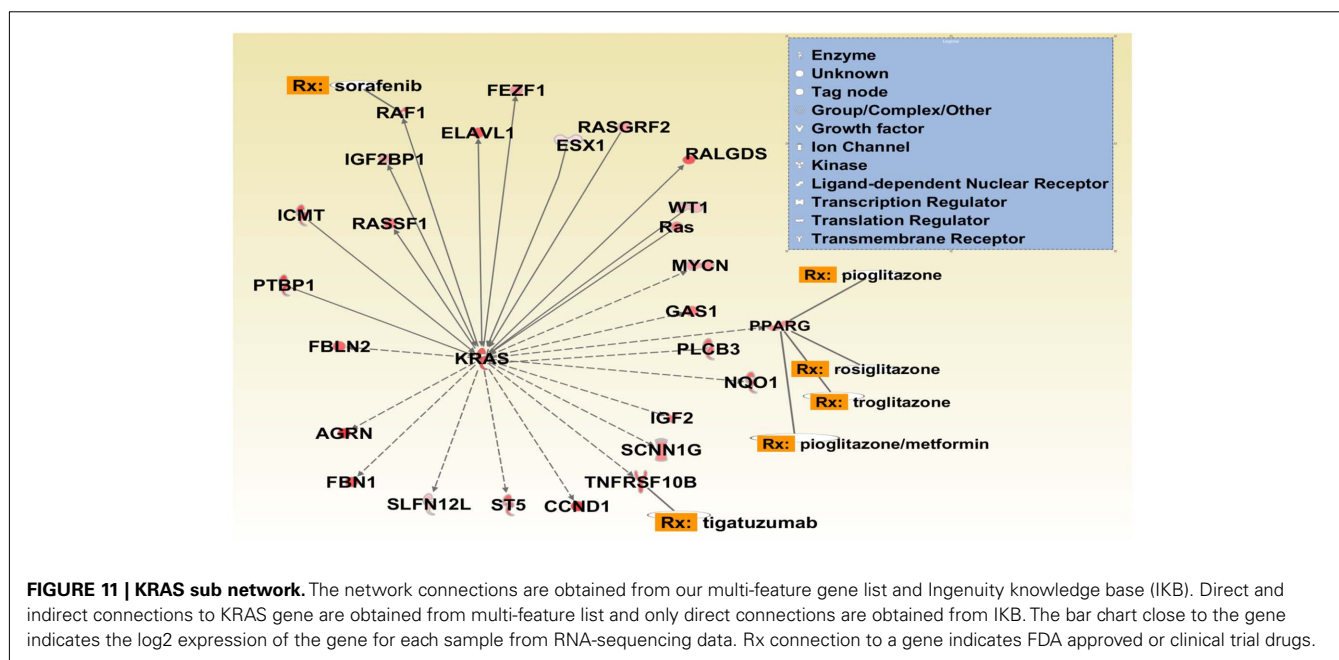


FIGURE 11 | KRAS sub network. The network connections are obtained from our multi-feature gene list and Ingenuity knowledge base (IKB). Direct and indirect connections to KRAS gene are obtained from multi-feature list and only direct connections are obtained from IKB. The bar chart close to the gene indicates the log2 expression of the gene for each sample from RNA-sequencing data. Rx connection to a gene indicates FDA approved or clinical trial drugs.

TNFR (Drosopoulos et al., 2005; Ji et al., 2006) and PPAR γ (Ignatenko et al., 2004) signaling pathways. Although our analysis is the first comprehensive transcriptome sequencing analysis concerned with the mutant *KRAS* phenotype, it does not have sufficient power to detect all genes associated to NSCLC tumors and *KRAS*. Thus, larger sample sizes will be desirable; and such analyses are ongoing at this time. However, we submit that the power of future studies is likely to be inherent in an integrated analysis, of the sort that we have defined here, which takes advantage of all of the genomic features that can be identified using RNA-Seq technology and leverages these technologies with new analytical tools. The tools that are available to quantify and integrate such features are currently in a state of rapid evolution as many investigators struggle with the complex issues related to building integrated systems models. The present model represents a first draft, rather than a rigorously defined genomic landscape model; and we anticipate that ongoing work in our laboratory and others will shortly lead to a better definition of the salient genomic features that distinguish *KRAS* mutational signaling events. However, our data demonstrate the utility of these approaches to identify critical druggable targets that can be

exploited to manage the very significant subset of NSCLC tumors with *KRAS* mutations.

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